

US 2006/0051371-A1  
Pinol *et al.*, Appl. No 10/535,416  
“Live attenuated vaccine against porcine  
pleuropneumonia”  
(HIPRA)

June 2009

# APP bacteria

Apx exotoxins (members of RTX toxins family):

- ApxI: strong haemolytic and high immunogenic  
Operon *apx/CABD* (*apx/C*, *apx/A*, *apx/B*, *apx/D* genes)

- ApxII: weak haemolytic and low immunogenic  
Operon *apx/ICΔB* (*apx/I/C*, *apx/I/A*, *apx/I/ΔB* genes)

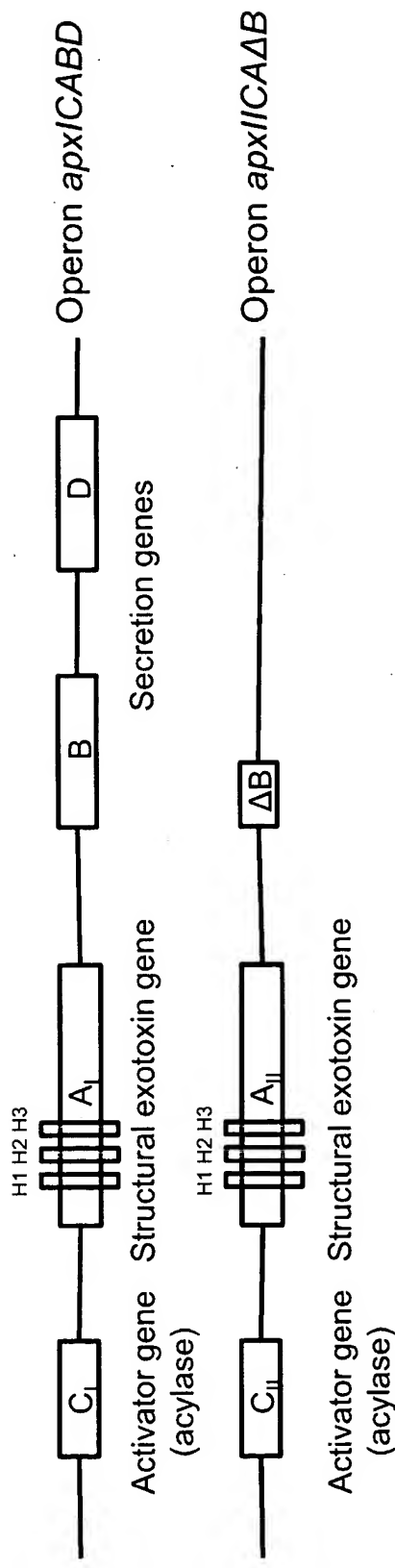
Genes:

- *apx/C*: activator gene for ApxI exotoxin
- *apx/A*: structural gene for ApxI exotoxin
- *apx/I/C*: activator gene for ApxII exotoxin
- *apx/I/A*: structural gene for ApxII exotoxin
- *apx/B* and *apx/D*: secretion genes of ApxI and ApxII exotoxins
- *apx/I/ΔB*: non-operative fragment

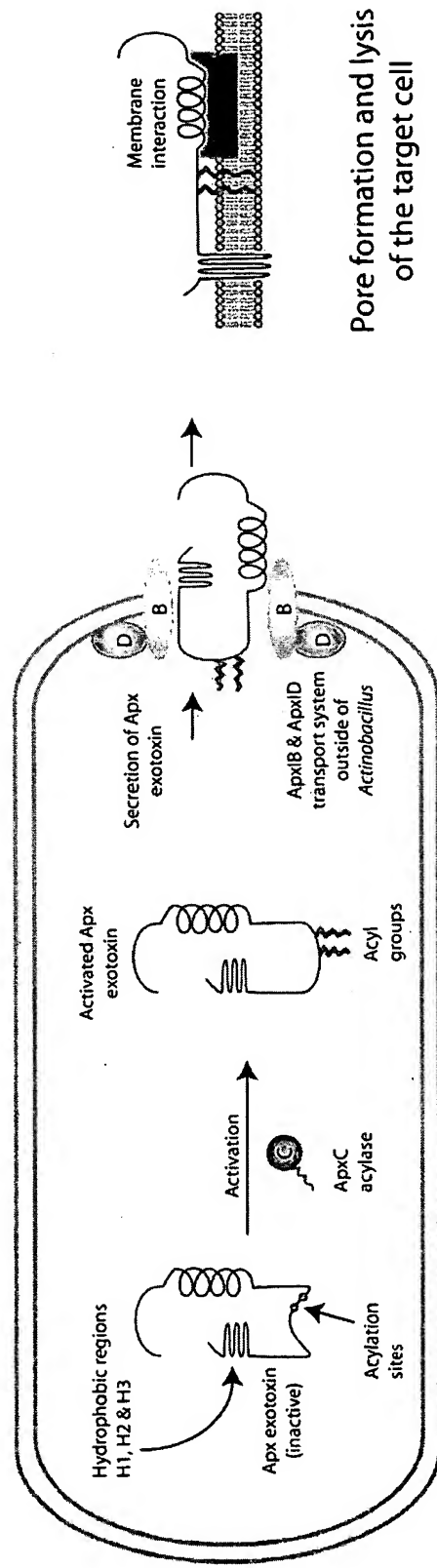
Apx exotoxins expression (several examples):

- Serotype 1: ApxI and ApxII exotoxins
- Serotype 10: only ApxI exotoxin
- Serotype 7: only ApxII exotoxin

## Structure of genes codifying ApxI $\alpha$ and ApxII $\alpha$ exotoxins

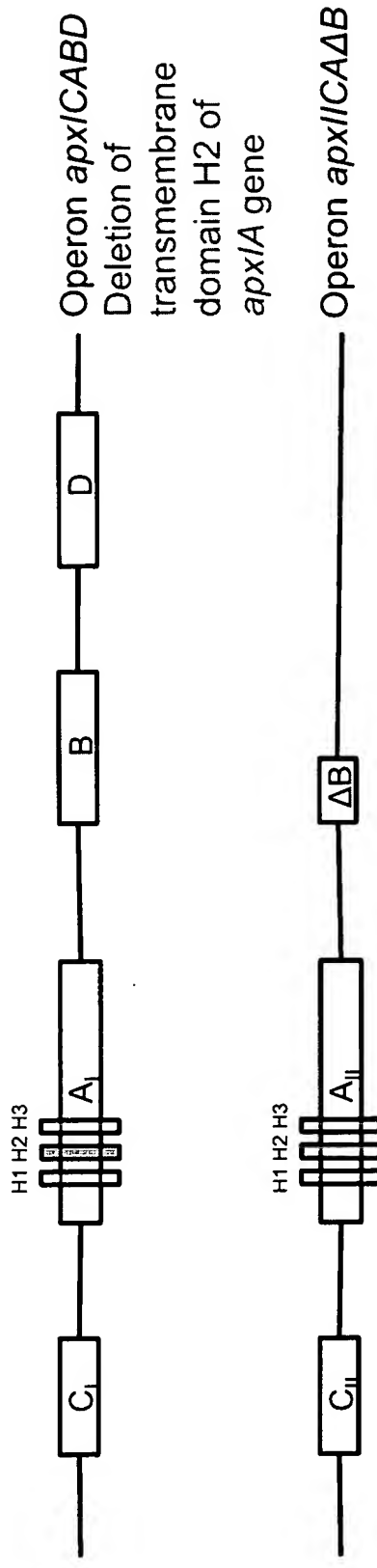


## Expression, activation and secretion of Apx exotoxins



# Pinol et al., US 2006/0051371-A1

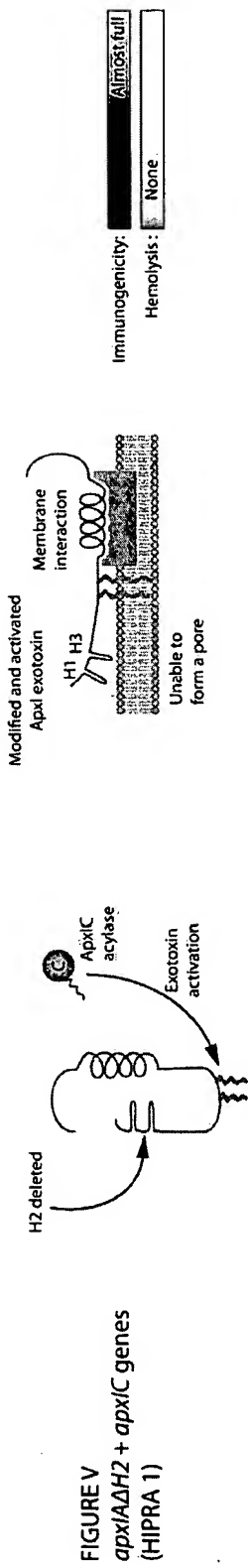
1) Deletion of a transmembrane domain of *apx/A* gen



Production and secretion of activated, but no haemolytic ApxI exotoxin, and activated ApxII exotoxin

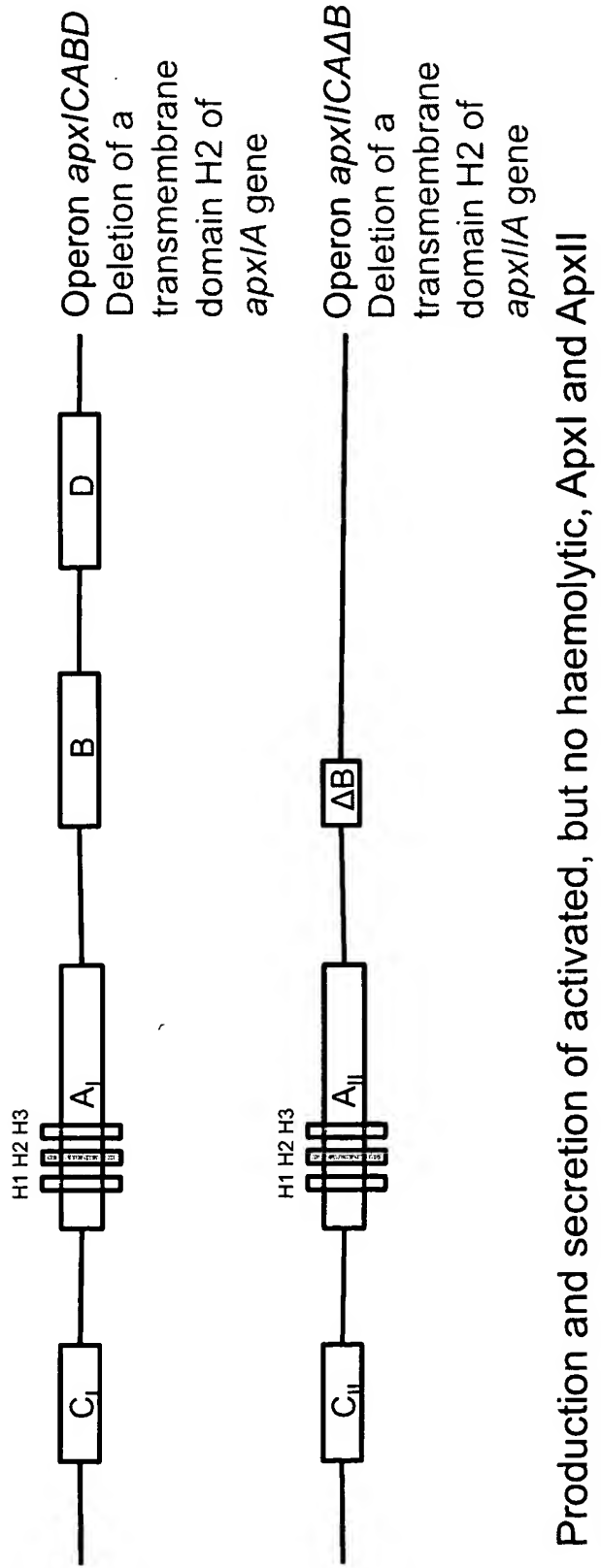
**High immunogenic** because ApxI and ApxII exotoxins are secreted

**Weak haemolytic** due to weak haemolytic activity of ApxII exotoxin  
(see Figure V)



# Pinol et al., US 2006/0051371-A1

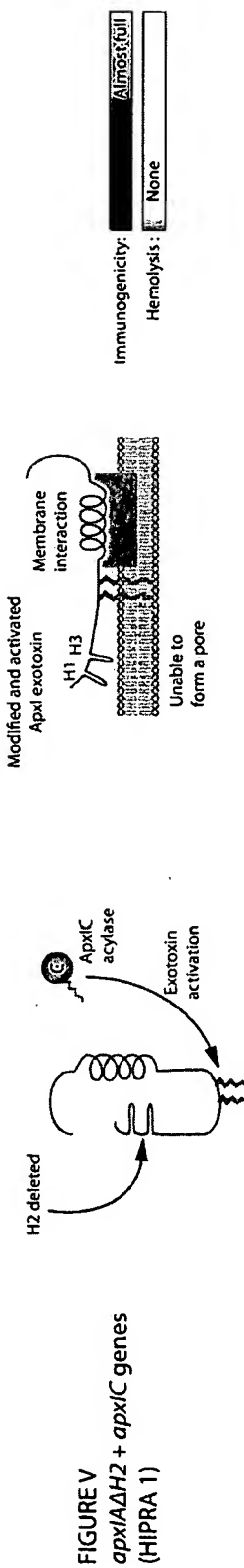
2) Deletion of a transmembrane domain of *apxIA* gene and of a transmembrane domain of *apxIIA* gene



Production and secretion of activated, but no haemolytic, ApxI and ApxII exotoxins

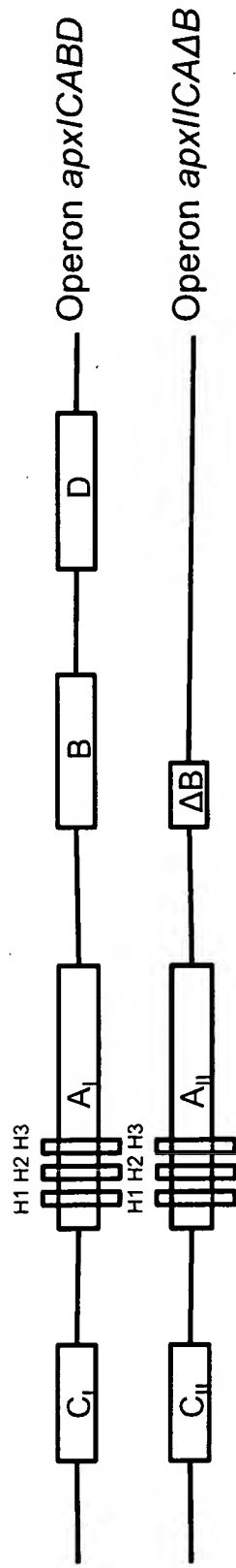
**High immunogenic** because ApxI and ApxII exotoxins are secreted

**Non-haemolytic** because modified ApxI and ApxII exotoxins are not capable to form pores (see Figure V)



Reimer *et al.*, Microbial Pathogenesis, 1995, 18: 197-209

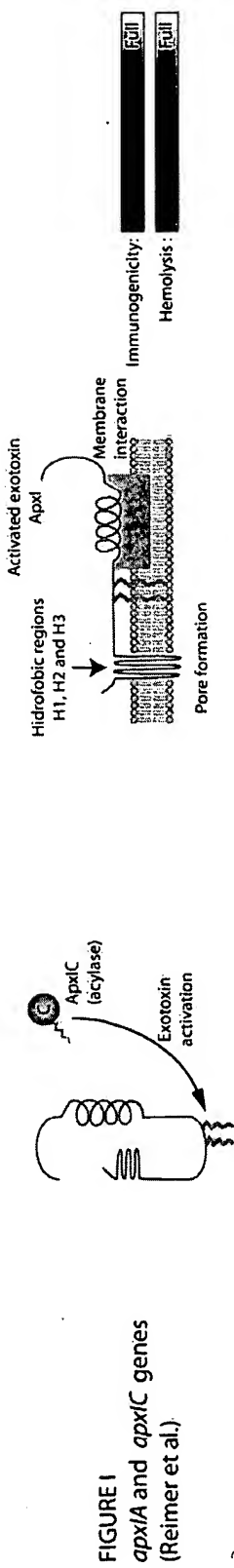
1) Strain J45: field isolate



Production and secretion of activated ApxI and ApxII exotoxins

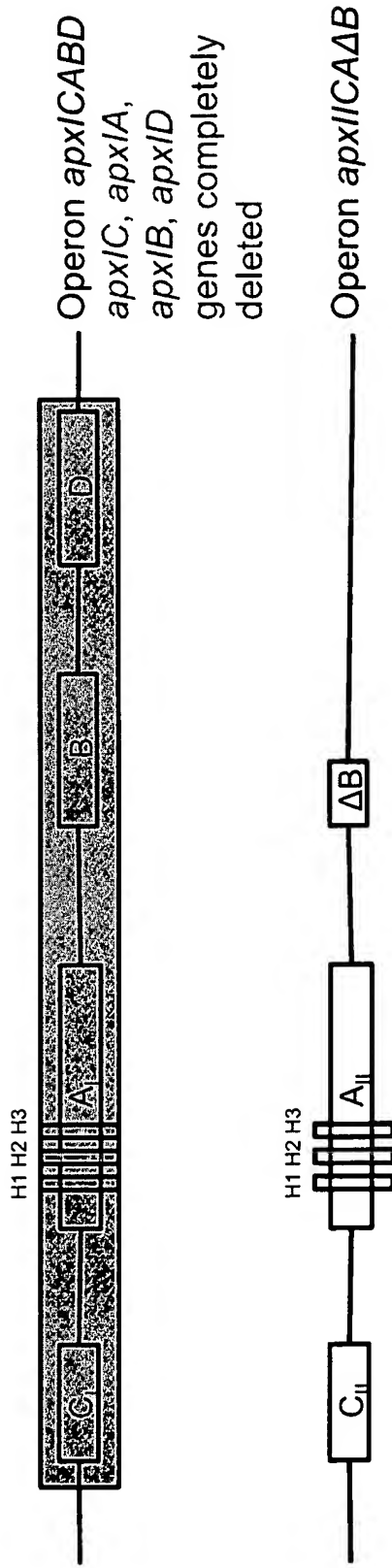
**High immunogenic** because it secretes ApxI and ApxII exotoxins

**Strong haemolytic** because ApxI and ApxII exotoxins are capable of forming pores  
(see Figure 1)



Reimer *et al.*, Microbial Pathogenesis, 1995, 18: 197-209

2) mlT4: chemical mutant



No production of ApxI exotoxin because of deletion of the whole *apx/CABD* operon

Production but no secretion of activated ApxII exotoxin because of deletion of *apx/B* and *apx/D* genes

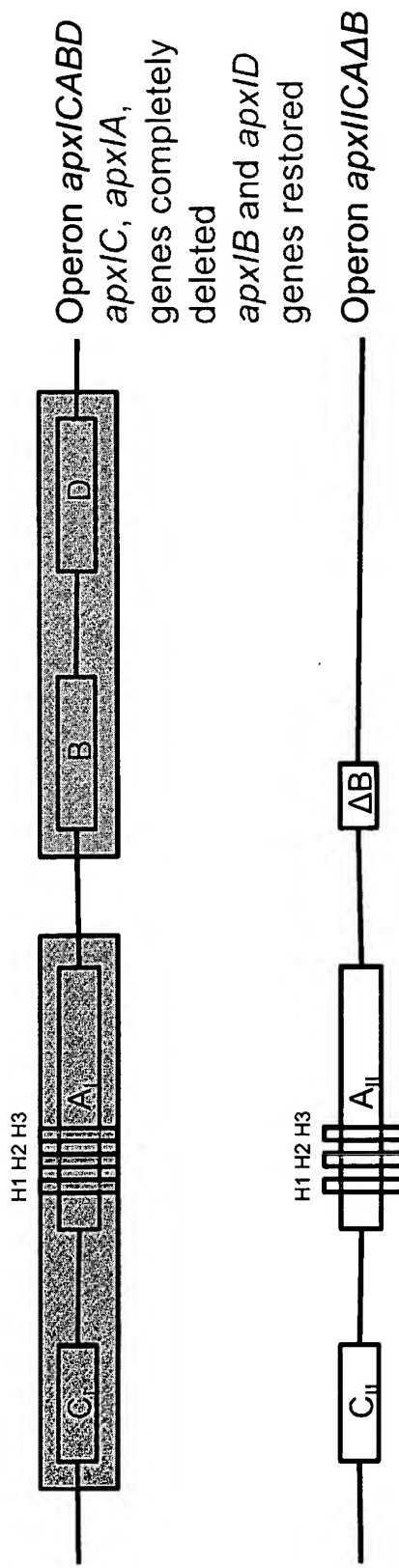
**Non-haemogenic** because ApxI and ApxII exotoxins are not secreted  
(see Figure II)



FIGURE II  
 $\Delta qpx/CABD$  genes  
(Reimer et al.)

Reimer *et al.*, Microbial Pathogenesis, 1995, 18: 197-209

3) Strain mIT4-H/pJFF801: chemical mutant with restored operon *apxIBD*



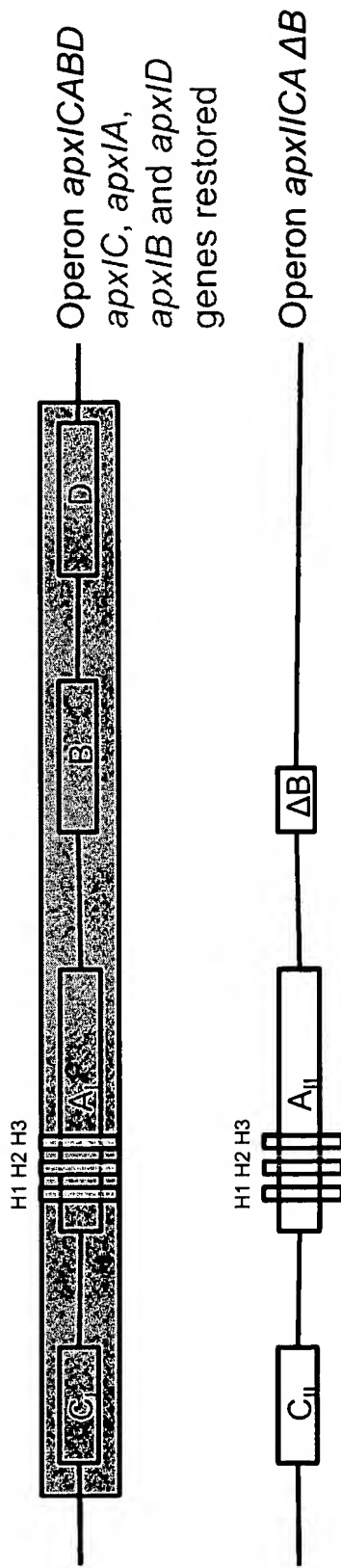
No production of activated ApxI exotoxin because *apxA* and *apxC* genes are completely deleted  
Production and secretion of activated ApxII exotoxin

**Low immunogenic** because ApxI exotoxin is not produced

**Weak haemolytic** because ApxII exotoxin is secreted

Reimer *et al.*, Microbial Pathogenesis, 1995, 18: 197-209

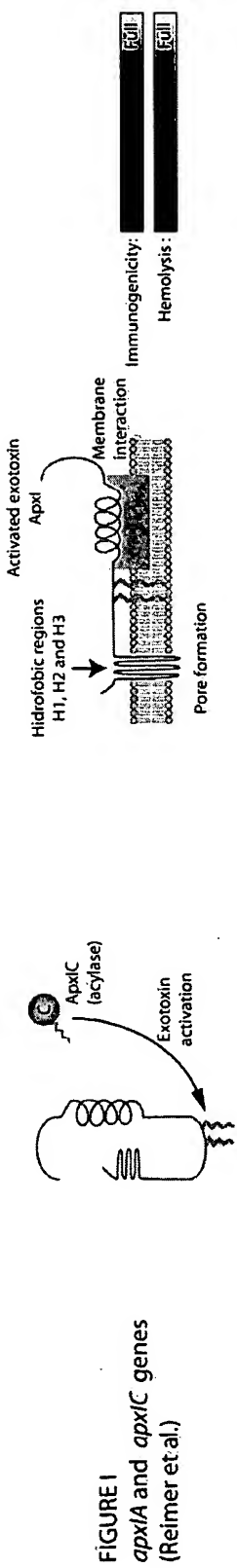
4) Strain mIT4-H/pJFF800: chemical mutant with restored operon *apx/CABD*



Production and secretion of activated ApxI and ApxII exotoxins

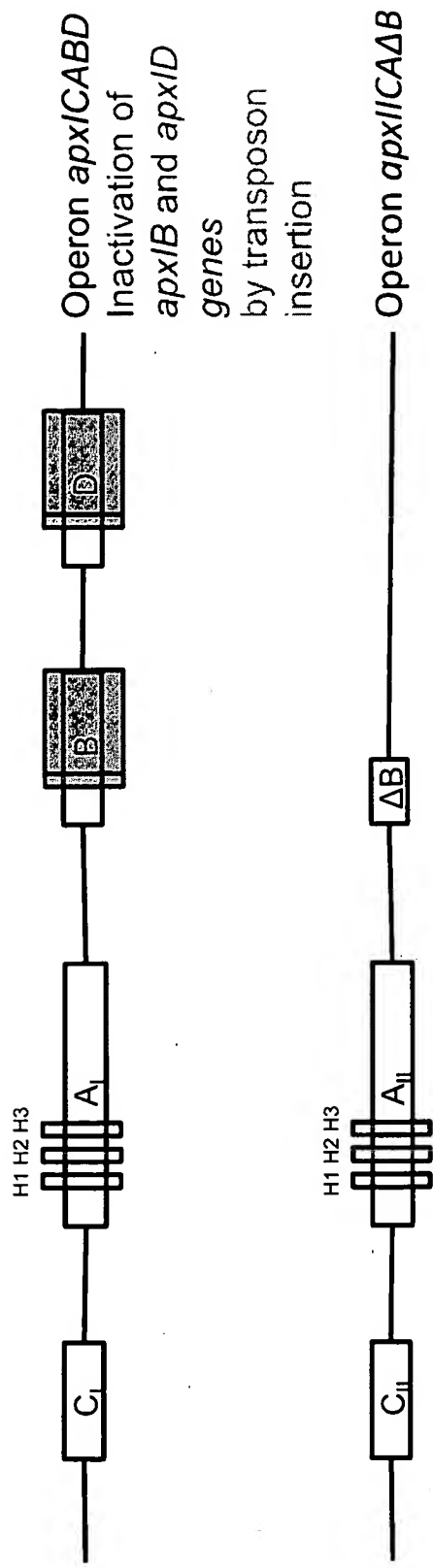
**High immunogenic** because it secretes ApxI and ApxII exotoxins

**Strong haemolytic** because ApxI and ApxII exotoxins are capable of forming pores  
(see Figure 1)



## MacInnes et al., US 6,019,984

Inactivation of *apxB* and *apxD* genes (secretion genes) by transposon insertion  
(Example 5)

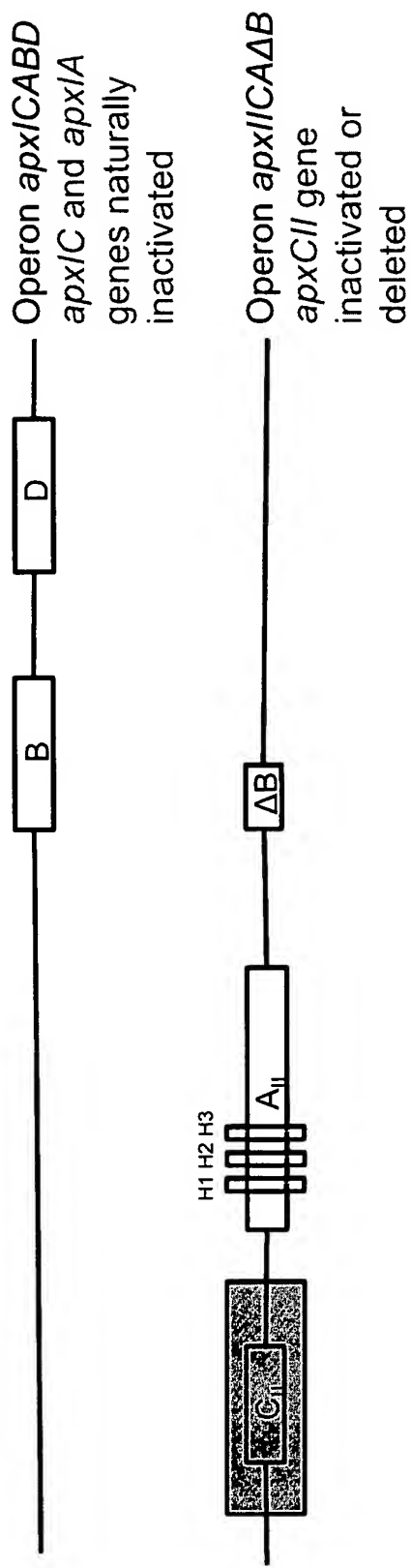


Production of cell-associated, activated ApxI and ApxII exotoxins, but they are not secreted  
(see Figure III)

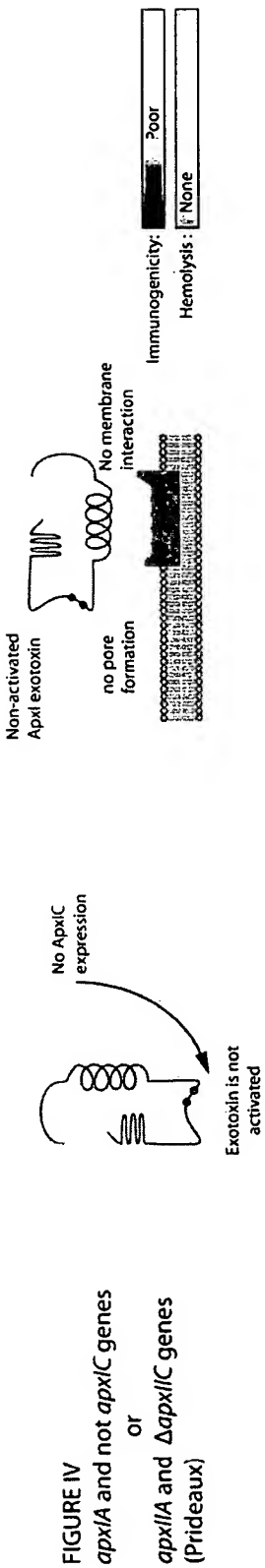


## Prideaux et al., US 6,472,183

- 1) Inactivation or deletion of *apxII/C* gene (activation gene) in wild strain HS93 (Serotype 7): strain HS93C- (Examples 10 and 11; column 20, lines 57-60; claims 1, 2, 3, 11, 12 and 14)

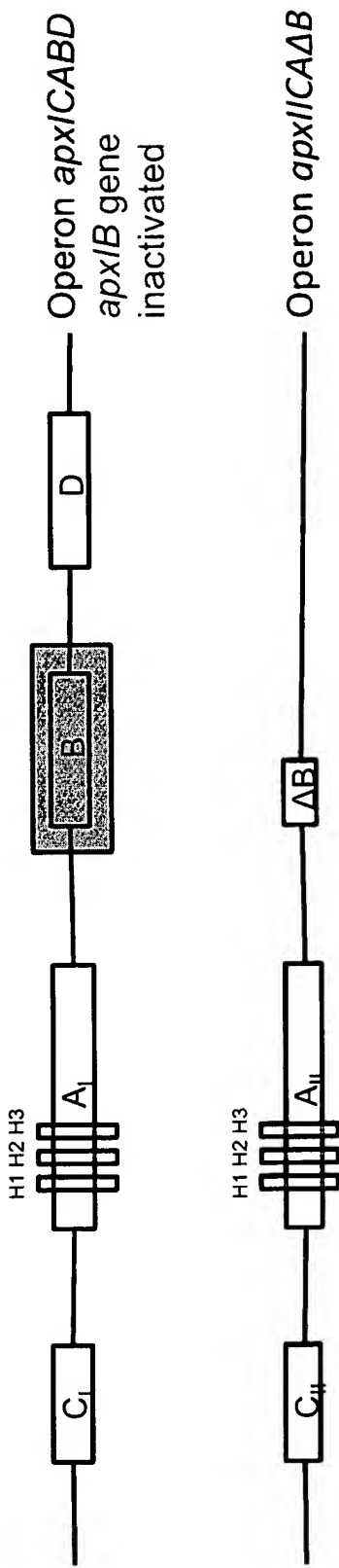


Production and secretion of non-activated ApxII exotoxin  
No production of ApxI exotoxin because of natural inactivation of  
*apxII/C* and *apxII/A* genes  
(see Figure IV)



Prideaux *et al.*, US 6,472,183

2) Inactivation of *apx/B* gene (secretion gene) in wild strain HS22 (Serovar 1):  
strain HS22B-  
(Examples 9 and 11; column 5, lines 21-24)

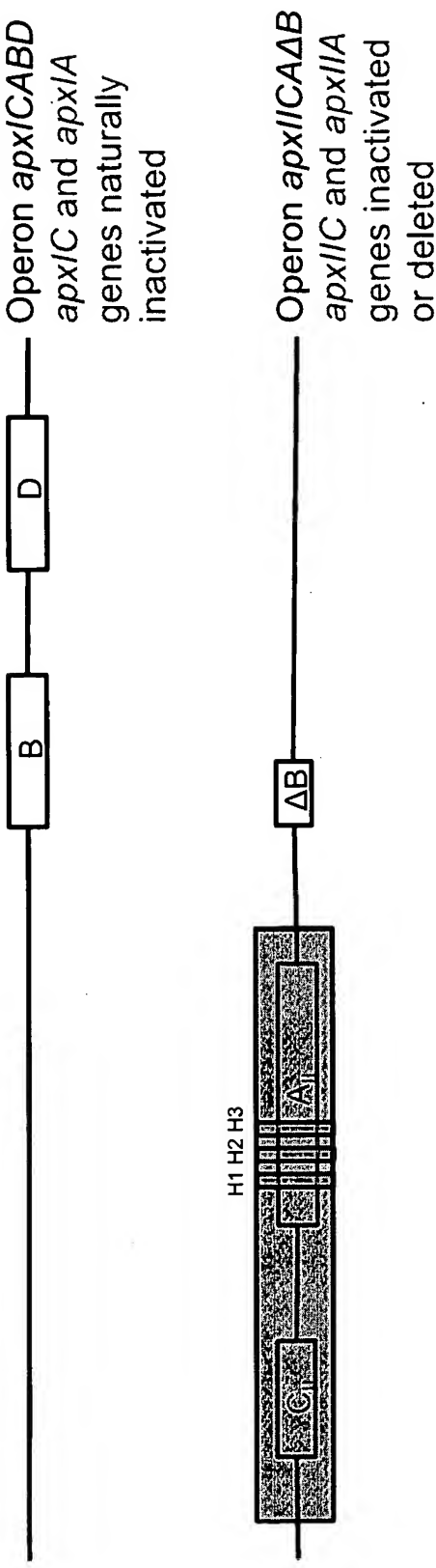


Production but no secretion of activated ApxI and ApxII exotoxins ,  
because of inactivation of *apx/B* gene  
(see Figure III)



## Prideaux *et al.*, US 6,472,183

3) Inactivation or deletion of *apxII/C* gene (activation gene) and *apxII/A* gene (structural exotoxin gene) of wild strain HS93 (Serovar 7): strain *Tox-*  
(Example 5)

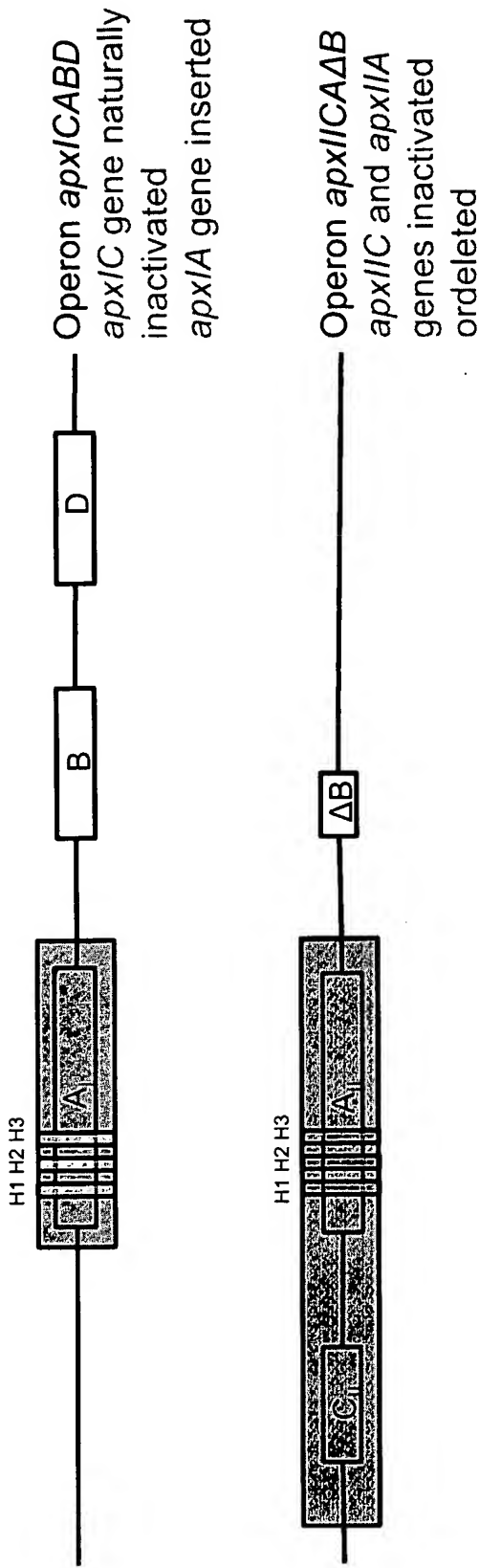


No secretion of exotoxins:

- ApxI exotoxin is naturally not produced
- ApxII exotoxin is not produced because of inactivation of *apxII/C* and *apxII/A* genes

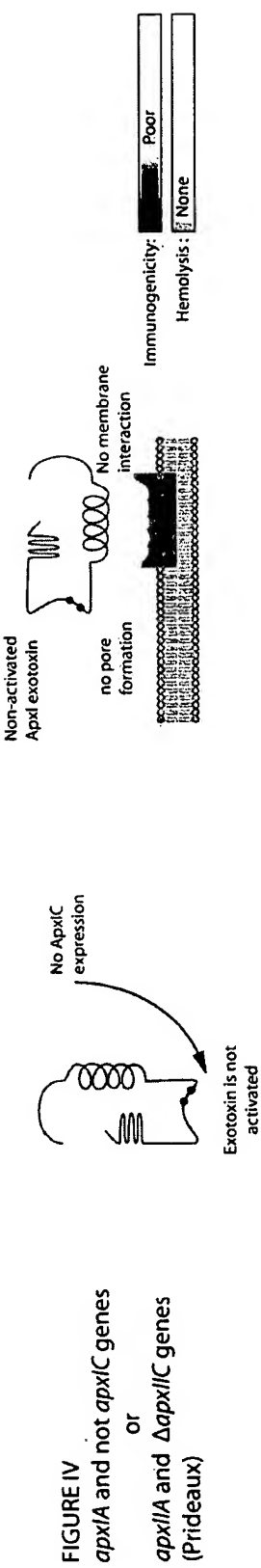
## Prideaux *et al.*, US 6,472,183

4) Insertion of *apx/A* gene (structural exotoxin gene) in strain Tox<sup>-</sup>:  
strain Tox<sup>-</sup>/pl63B-TIK  
(Examples 3,4, 5 and 6; column 4, lines 58-65)



Production of non-activated ApxI exotoxin because *apx/C* gene is naturally inactivated

No production of ApxII exotoxin because *apx/C* and *apx/A* genes are inactivated  
(see Figure IV)



# Conclusions

Pinol *et al.*, US 2006/0051371-A1

**1.- Technical concept**

Mutation (deletion) in a transmembrane domain of exotoxin A genes

**2.- Novelty**

None of the documents of the state of the art discloses a mutation (deletion) in a transmembrane domain of *apxI/A* gene, with or without a mutation (deletion) in a transmembrane domain of *apxII/A* gene.

**3.- Inventive step**

Once a mutation (deletion) in a transmembrane domain of *apxI/A* gene or in a transmembrane domains of *apxI/A* and *apxII/A* genes has been performed, it would not have been obvious for the skilled person that the protein:

- a) would maintain the structure
- b) would be secreted
- c) would not be haemolytic
- d) would be immunogenic
- e) would be immunoprotective

**Pinol et al., US 2006/0051371-A1**

4.- Applicant strains are highly immunogenic and non-haemolytic because:

- a) they produce and secrete activated ApxI and ApxII exotoxins
- b) these exotoxins are not capable of forming pores

5.- So, a mutation (deletion) carried out in a transmembrane domain of the *apxIA* gene, with or without a mutation (deletion) in a transmembrane domain of the *apxIIA* gene surprisingly resulted in:

- maintenance of the structure of ApxI and ApxII exotoxins,
- secretion of the ApxI and ApxII exotoxins,
- non-haemolytic activity,
- immunogenicity and
- immunoprotective characteristics

**Pinol et al., US 2006/0051371-A1**

- 6.- Claims 13, 14, 15, 16, 17 and 19 currently on file are drawn to immunogenic, non-haemolytic APP strains comprising at least a mutation (deletion) in a transmembrane domain region of the *apx/A* gene, and optionally, a mutation in a transmembrane domain region of the *apx//A* gene.
- 7.- Any of the documents cited in the prior art do not disclose, suggest or teach APP strains obtained by mutation (deletion) in a segment of the transmembrane domain region of the *apx/A* gene, with or without a mutation (deletion) in a segment of the transmembrane domain region of the *apx//A* gene.
- 8.- All documents cited in the prior art were driven by the same idea and purpose: that the absence of the main virulence factor of APP, i.e. Apx toxins, (by deletion, or non-activation, or non-secretion) would result in a non-virulent (non-haemolytic), but protective strain.

**Pinol et al., US 2006/0051371-A1**

- 9.- In APP this strategy resulted less efficient than in other microorganisms, because Apx toxins need to be activated and secreted in order to induce a high level of immunoprotection.
- 10.- It would not have been obvious for the skilled person that a mutation (deletion) in a transmembrane domain region of the *apx/A* gene, with or without a mutation (deletion) in a transmembrane domain region of the *apxII/A* gene would lead to an APP strain expressing and secreting activated Apx toxins, so maintaining its immunogenic properties, but not its haemolytic activity, resulting consequently in a non-virulent strain being not capable of producing pores in target cells.

# **Illustrated summary with idealized structures and mechanisms**

(without being bound to the theory)

